PATENT COOPERATION TREATY INTERNATIONAL PRELIMINARY EXAMINATION REPORT

PCT/GE91/00212 Receiving Office UK PATENT OFFICE Applicant (Name) AMERSHAM INTERNATIONAL PLC ET AL BASIS OF	International Filing Date 13 February 1991 (13.02.91) Priority Date Claimed 13 February 1990 (13.02.90)	
UK PATENT OFFICE Applicant (Name) AMERSHAM INTERNATIONAL PLC ET AL BASIS OF	Priority Date Claimed 13 February 1990 (13.02.90)	
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AMERSHAM INTERNATIONAL PLC ET AL BASIS OF		
AMERSHAM INTERNATIONAL PLC ET AL BASIS OF	REPORT	
BASIS OF	REPORT	
AMENDMENTS AND/OR RECTIFICATIONS!* — The amendments an	REPORT	
AMENDMENTS AND/OR RECTIFICATIONS! - The amendments an		
Authority in respect of the claims, the description, and/or drawings in the	d/or rectifications made before this International Preliminary Exame above-identified international application are annexed to this rep	nininç ort.
. X This report has been established on the basis of the following	application documents:	
X the application documents as filed		
description, pages	as originally filed	
description, pages	filed with your letter of	
description, pages	filed with your letter of	
description, pages	filed with your letter of	
claim(s)	as originally filed	
claim(s)	filed with your letter of	
claim(s)	filed with your letter of	
claim(s)	filed with your letter of	
drawings, sheet/fig.	as originally filed	
drawings, sheet/fig.	filed with your letter of	
. The amendments resulted in the cancellation of the following sheet	B:	· • • • ·
This report has been established as if the amendments indicated on	the extra sheet have not been made, since, for the reasons indicated	J. the
have been considered to go beyond the disclosure as filed.	•	
. PRIORITY 2		
	·	
a. This report has been established as if no priority has been cla requested:	imed due to the failure to furnish within the prescribed time li	nit th
copy of the earlier application whose priority has been claim	med.	
translation of the earlier application whose priority has been	n claimed.	
b. This report has been established as if no priority has been c	aimed due to the fact that the priority claim has been found i	nvali
Thus, for the purposes of this report, the international filing date indic	ated above is considered to be the relevant date.	
 Where replacement sheets are annexed to this report, a translation of the limit applicable under PCT Article 39(1). 	se replacement sheets must be furnished to the elected Offices within	the ti

	BASIS OF REPORT (Continued)
3.	UNITY OF INVENTION 3 — The international application does not comply with the requirement of unity of invention.
	a. In response to an invitation to restrict or pay additional fees the applicant has:
	restricted the claims.
	paid additional fees.
	paid additional fees under protest. Where requested by the applicant, the text of the protest together with the decision taken thereon are annexed to this report.
	neither restricted nor paid additional fees.
	b. No invitation has been issued. The opinion of this International Preliminary Examining Authority is that the international application does not comply with the requirement of unity of invention for the following reasons. (specify)
	\cdot
	 Consequently, the following parts of the international application were the subject of international preliminary examination in establishing this report:
	all parts.
	the parts relating to the restricted claims, that is claims Nos.
	the parts relating to the main invention, that is claims Nos
•.	NON-ESTABLISHMENT OF REPORT ON QUESTIONS OF NOVELTY, INVENTIVE STEP OR INDUSTRIAL APPLICABILITY 4 The questions of whether the claimed invention appears to be novel, to involve an inventive step or to be industrially applicable
	have not for the reasons indicated been gone into in respect of:
	a. the entire international application
	b. claims Nos.
	for the following reasons:
	Said international application, or said claims Nos relate to the following subject matter which does not require an international preliminary examination. (specify)
	The description, claims, or drawings (indicate particular elements) or said claims Nos are so unclear that no meaningful opinion could be formed.
	The claims, or said claims Nos are so inadequately supported by the description that no meaningful opinion could be formed.
	Said claims Nos are dependent claims and are not drafted in accordance with the second and third sentences of PCT
	Rule 6.4(a).
	\cdot

CLASSIFICATION OF SUBJECT MATTER (If several classification symbols apply, indicate all.) 5

According to International Patent Classification (IPC) or to both National Classification and IPC

 IPC^{5} B03C 1/00: C07K 3/24: 3/28: C12N 1/02. 7/02. 5/00

REASONED STATEMENT AS TO CLAIMS MEETING CRITERIA OF NOVELTY (N), INVENTIVE STEP (IS) AND INDUSTRIAL APPLICABILITY (IA) AND CITATIONS AND EXPLANATIONS SUPPORTING SUCH STATEMENT

		SUPPORTING SUCH STATEMENT
CLAIM NUMBER	STATEMENT (CRITERIA)	CITATIONS AND EXPLANATIONS
1-12	YES (N. IS. IA)	All claims meet the requirements of industrial applicability novelty and inventive step.
·		They are distinguished from the documents cited by the International Searching Authority in that there is no disclosure of the suspension of magnetic particles in a polymer solution prior to precipitating the polymer out of the solution.
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	NON-WRITTER	DISCLOSURES .	
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	m or contents of the international		ATION "
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Form PCT/IPEA/409 (last sheet) (January 1985)

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See notes on accompanying sheet

INTERNATIONAL APPLICATION UNDER THE PATENT COOPERATION TREATY

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INTERNAT	TONAL TE:	13 Fe	bruary 1991 - Z-9)	
(Stamp) Name of rec	PCT Inte	rnational	atent Office Application ational Application	
Applicant's (indicated by	or agent's file refer applicant if desi	rence red)	PP/8620	

TATEM COOPERATION TREATY	FILING DATE: 13—21—9)
REQUEST THE UNDERSIGNED REQUESTS THAT THE PRESENT INTERNATIONAL APPLICATION BE PROCESSED ACCORDING TO THE PATENT COOPERATION TREATY	United Kingdom Patent Office PCT International Application Name of receiving Office and "PCT International Application"
TO THE PARENT COOPERATION TREAT	Applicant's or agent's file reference (indicated by applicant if desired) PP/8620
Box No. 1 TITLE OF INVENTION	
PRECIPITATING POLYMERS	
(includes, where applicable, a legal entity) is involved, continue	
The person identified in this box is (mark one check-box only):	applicant and applicant inventor*
Name and address: •• AMERSHAM INTERNATIONAL Amersham Place Little Chalfont	PLC
Buckinghamshire HP7 9NA	
ENGLAND G.B.	
Telephone number (including area code): Telegraphic address	Teleprinter address:
State of nationality: "C.B"	State of residence: * CG.B
The person identified in this box is applicant for the purposes of	(mark one check-box only):
all designated States except the United States of America	the United States the States indicated in the "Supplemental Box"
where applicable, a legal entity). If the following two sub-boxes	HER) INVENTORS. IF ANY: DESIGNATED STATES FOR separate sub-box has to be filled in in respect of each person uncludes, are insufficient, continue in the "Supplemental Box." (giving there for in the following two sub-boxes) or by using a "continuation sheet." [applicant and applicant inventor only only only only only only only only
If the person identified in this sub-box is applicant (or applicant of	and inventor), indicate also:
State of nationality: 4.B	State of residence:
and whether that person is applicant for the purposes of (mark o	ne check-box only): the United States the States indicated
States the United States of America	of America only in the "Supplemental Box"
The person identified in this sub-box is (mark one check-box only Name and address: **	y): applicant and applicant inventor only
If the person identified in this sub-box is applicant for applicant of	
State of nationality: and whether that person is applicant for the purposes of (mark or	State of residence: *
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and the me measure and indications in the Subbishells in	y name first followed by the given name(s). Indicate the name of a legal

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	PENNANT, PYERS				
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	Chancery Lane				
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•	5-8393			071-430-2262	
	DESIGNATION OF GROUPS OF STATES	OP ST	ATE		CINDS OF
ROTECT	ION OR TREATMENT. The following designation	s are here	by m	ide (please mark the applicable check-	boxes):
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<u>X</u> EP	European Patent ⁽²⁾ : AT Austria, BE Belgium, DK Denmark, ES Spain, FR France, GB Unit NL Netherlands, SE Sweden, and any other State which is a Contracting State of the	ed Kingo	dom.	GR Greece, IT Italy, LU Luxemb	Germany, courg,
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Patent Office (see also the "Notes to Box No. V").

(3) If another kind of protection or a title of addition or, in the United States of America, treats in-part is desired, specify according to the instructions given in the "Notes to Box No. V."

R 1B CHANGE OF ADDRESS SEE #12

No. VI PRIORITY CI	AIM (IF ANY). The priority of the	following earlier application(s) is	hereby claimed:
Country (country in which it was filed if national applica- tion; one of the countries for which it was filed if regional or international application)	Filing Date (day. month, year)	Application No.	Office of filing (fill in only if the earlier application is an international application or a regional application)
□ G.B.	13 February 1990	90 03 253.3	
(2)	13-2-1990		
(3)			
When the earlier application w Office, the applicant may, agai	dicate country and/or Office of filing as filed with the Office which, for the nst payment of the required fee, ask the reby requested to prepare and transmy of the earlier applications identified a	purposes of the present internation following: it to the International Bureau a	certified copy of the above-men-
Searching Authority has alread to the extent possible, on the re	ARCH (IF ANY). Fill in where a sea y been requested (or completed) and the sults of the said earlier search. Identified or by reference to the search request.	ne said Authority is now requested y such search or request either by	to base the international search.
International application number and country for region Office) of other application:		International/regional/nationa filing date:	13 February 1990 .
Date of request for search:	27 February 1990	Number (if available) given to search request:	RS 85977 GB
PENNANT, PYER If the present Request form is signed by the applicant is requested. A copy thereof must be	signed on behalf of any applicant by ired. If in such case it is desired to ma	v an agent, a separate power of a like use of a general power of atto	ttomey appointing the agent and mey (deposited with the receiving
Box No. IX CHECK LIST	(To be filled in by the Applicant)	This international application items marked below:	n as filed is accompanied by the
This international application of sheets:	tion contains the following number	1. X separate signed power	of attorney
1 request3	sheets	2. Copy of general power	of attorney
2. description3	sheets sheets	3. priority document(s) (see Box No. VI)
4. abstract1	sheets	4. receipt of the fees pai	d or revenue stamps
5. drawings4	Tetal 27 sheets	5. X cheque for the payme	
	Total 2/ sheets	6. request to charge dep	
Figure numberis suggested to accompan	y the abstract for publication.		Patents Form 24/77, Earlier search report.
Date of actual receipt of	(The following is to be filler the purported international application	17 Fahrmann	1991 13-2-91
	receipt due to later but timely received the purported international application		
3. Date of timely receipt of	the required corrections under Article	11 of the PCT:	•
4. Drawings Received	No Drawings		
Date of receipt of the record		1991 (0 5. 03. 91)	

Form PCT/RO/101 (last short) (January 1991)

PATENT COOPERATION TREATY INTERNATIONAL SEARCH REPORT

ADDITION OF INTERNATIONAL ARRIVATION		Applicant's or Agent's File Reference
IDENTIFICATION OF INTERNATIONAL APPLICATION		PP/8620
International Application No.	International	Filing Date
PCT/GB 91/00212		13th February 1991
Receiving Office	Priority Date	Claimed
RO/GB		13th February 1990
Applicant		
AMERSHAM INTERNATIONAL PLC e		
I. CERTAIN CLAIMS WERE FOUND UNSEARCHABLE 1		
II. UNITY OF INVENTION IS LACKING 2 (Observations	on supplement	ai sheet (2))
III. TITLE, ABSTRACT AND FIGURE OF DRAWING		
Title. X Abstract. Abstract.	oplicant: 3	
2. The texts established by this International Searching Authority of (Title. Abstract.	the following Ind	sicated Items are set forth below:
Method to isolate macromolectatractable beads which do no macromolecules.		
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The sale of the state of the sale of the s		
Text of the abstract continued on supplemental sheet (1) 3. a. The definitive contents of the abstract are established by the	is International	Searching Authority se aronneed in form DCT/ISA/2004
b. This report is incomplete as far as the abstract is concerned by this international Searching Authority has not expired.	d as the time lim	
4. Figure to be published with the abstract 6	٠	
Figure No. 1a. None of the figures		
as suggested by the applicant		
because the applicant failed to suggest a figure		
because this figure better characterizes the invention		1 4

INTERNATIONAL SEARCH REPORT

International Application No PCT/GB 91/00212

	IFICATION													ii) ⁶				
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111. DO	CUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET	-)
Category *	Citation of Document, 11 with indication, where appropriate, of the relevant passages	Relevant to Claim No.
A	WO, A, 8401503 (COULTER et al.) 26 April 1984 see the whole document; especially page 7, lines 1-5	1-12
A	EP, A, 0281390 (LYLE et al.) 7 September 1988 see examples	1-12
A	Chemical Abstracts, vol. 112, no. 11, 12 March 1990, (Columbus, Ohio, US), S. Flygare et al.: "Magnetic aqueous two-phase separation in preparative applications", see page 606, abstract 96880s,	1-12
	& Enzyme Microb. Technol. 1990, 12(2), 95-103	1 10
A	Chemical Abstracts, vol. 82, no. 18, 5 May 1975, (Columbus, Ohio, US), G. Bitton et al.: "Removal of Escherichia coli bacteriophage T7 by magnetic filtration", see page 278, abstract 115953y, & Water Res. 1974, 8(8), 549-51	1-12
A	Chemical Abstracts, vol. 77, no. 14, 2 October 1972, (Columbus, Ohio, US), J. Warren: "New purification procedure for biological vaccines (adsorption on magnetic iron oxides)", see page 308, abstract 92767w, & Immunization Jap. Encephalitis, Conf. 1969 (Pub. 1971), 152-4	1-12
A	Chemical Abstracts, vol. 95, no. 9, 31 August 1981, (Columbus, Ohio, US), P.A. Munro et al.: "Magnetic seeding to aid recovery of biologica precipitates", see pages 394-395, abstract 76474q, & Biotechnol. Lett. 1981, 3(6), 297-302	1-12
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

GB 9100212

SA 44598

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on 26/06/91

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Patent document cited in search report	Publication date	Patent family member(s)	Publication date
EP-A- 0162819	27-11-85	AU-A- 3916985 JP-A- 60210766 SE-A- 8401125	23-10-85
US-A- 4001197	04-01-77	JP-C- 942629 JP-A- 51150779 JP-B- 53019821	24-12-76
US-A- 3470067	30-09-69	None	
BE-A- 686243	28-02-67	None	
WO-A- 8401503	26-04-84	US-A- 4508625 AU-A- 2205683 DE-T- 3390261 EP-A- 0124579 JP-T- 59501867 SE-B- 452258 SE-A- 8403217 CA-A- 1228053	3 04-05-84 1 10-01-85 1 14-11-84 7 08-11-84 3 23-11-87 7 15-06-84
EP-A- 0281390	07-09-88	AU-A- 1426988 JP-T- 1502319 WO-A- 8806633	3 17-08-89

	17 Rec'd PCI/PTO 1 APR 1991
PATENT COOPERATION TREATY	
	international application no. pct/gb91/00212
NOTIFICATION TO THE DESIGNATED OFFICE OF RECEIPT OF RECORD COPY issued under PCT Rule 24.2(a)	To: United States Patent and Trademark Office Washington, D.C.
	in its capacity as a designated Office
	From: The International Bureau of WIPO 1211 Geneva 20 Switzerland
NAME(S) OF APPLICANT(S):	
REEVE, Michael, Alan	
INTERNATIONAL FILING DATE:	February 1991 (13.02.91)
PRIORITY DATE(S) CLAIMED:	February 1990 (13.02.90)
DATE OF RECEIPT OF RECORD COPY 1	BY INTERNATIONAL BUREAU: March 1991 (05.03.91)
	D. Collier (Authorized Officer)

Form PCT/IB/302 (January 1984)

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United States Patent and Trademark Office Washington, D.C. the INTERNATIONAL BUREAU of the WORLD INTELLECTUAL PROPERTY ORGANIZATION

NOTIFICATION CONCERNING DOCUMENTS TRANSMITTED

Issued pursuant to PCT Article 36(3)(a)

(as elected Office)

Date of Mailing:

20 February 1992 (20.02.92)

NOTIFICATION

The International Bureau transmits herewith the following documents and number thereof:

1 (number of) copy(s) of the international preliminary examination report (Article 36(3)(a)).

This notification is sent to the above addressee in its capacity as an elected Office.

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Switzerland

<u>Authorised Officer:</u>

M. Abidine



International Application No PCT/GB 91/00212

			memanent Application to 2 3 2 7				
I. CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) ⁶							
According		onal Patent Classification (IPC) or to both Natio					
IPC ⁵	B 0	3 C 1/00, C 07 K 3/24,	3/28, C 12 N 1/02,	, 7/02 , 5/00			
1PC :							
II. FIELDS	SEARCH	IED					
		Minimum Documen	tation Searched 7				
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IPC ⁵		в 03 с, с 07 к, с	12 N				
IPC		B 03 C, C 07 R, C	IZ N				
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		to the Extent that such Documents	are Included in the Fields Searched				
III. DOCU		ONSIDERED TO BE RELEVANT		I a v v v v v			
Category *	Citat	ion of Document, 11 with Indication, where appr	opriate, of the relevant passages 12	Relevant to Claim No. 13			
X	EP,	A, 0162819 (PERO et a	1.)	1-4,6-8			
İ		27 November 1985	•				
		see the whole documen	t; in particular				
		page 2, line 25 - pag	e 4. line 31				
			,				
х	IIS.	A, 4001197 (MITCHELL	et al.)	1-4.6-8.12			
1	00,	4 January 1977	cc a1.,	1 4,0 0,12			
			+. ogpogiallæ				
		see the whole documen		1			
		figure 1; columns 6-8	, claims				
							
X	US,	A, 3470067 (WARREN et	al.)	1-4,6-8			
		30 September 1969					
		see the whole documen	t; especially	1			
		columns 1,2, summary;	column 8, claims				
			•				
х	BE.	A, 686243 (PFIZER) 28	February 1967	1-4,6-8			
	,	see the whole documen		1 2 1,0 0			
		pages 7-8, examples 1		1			
		claims	,2, pages 14-13,				
		Claims					
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	-	s of cited documents: 10	"T" later document published after to or priority date and not in confi				
		ning the general state of the art which is not be of particular relevance	cited to understand the princip	e or theory underlying the			
		nt but published on or after the international	invention "X" document of particular relevan	set the claimed invention			
តិដែក	g date		cannot be considered novel or				
"L" doc	ument which is cited	th may throw doubts on priority claim(s) or to establish the publication date of another	involve an inventive step "Y" document of particular relevan	see the claimed invention			
cita	tion or othe	er special resson (as specified)	cannot be considered to involve	an inventive step when the			
	ument refe er means	rring to an oral disclosure, use, exhibition or	document is combined with one ments, such combination being				
"P" doc	ument pub	ished prior to the international filing date but	in the art.				
		priority date claimed	"A" document member of the same	patent family			
IV. CERT	IFICATIO	M.					
Date of the	• Actual Co	empletion of the International Search	Date of Mailing of this International S	earch Report			
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Internation	nai Searchir	ng Authority	Signature of Authorized Officer				
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111. DO	CUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET	Γ)
Category *	Citation of Document, 13 with Indication, where appropriate, of the relevant passages	Relevant to Claim No.
A	WO, A, 8401503 (COULTER et al.) 26 April 1984 see the whole document; especially page 7, lines 1-5	1-12
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO.

GB 9100212 SA 44598

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Patent document cited in search report	Publication date	Patent fa member		Publication date
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US-A- 3470067	30-09-69	None		
BE-A- 686243	28-02-67	None		
WO-A- 8401503	26-04-84	AU-A- DE-T- EP-A- JP-T- 5 SE-B- SE-A-	4508625 2205683 3390261 0124579 9501867 452258 8403217 1228053	02-04-85 04-05-84 10-01-85 14-11-84 08-11-84 23-11-87 15-06-84 13-10-87
EP-A- 0281390	07-09-88	JP-T-	1426988 1502319 8806633	26-09-88 17-08-89 07-09-88

REC'D 19 FEB 1992

PATENT COOPERATION TREATY INTERNATIONAL PRELIMINARY EXAMINATION REPORT

DENTIFICATION OF THE INTERNATIONAL APPLICATION	Applicant's or Agent's File Reference PP/8620					
nternational Application No.	International Filing Date					
PCT/GE91/00212	13 February 1991 (13.02.91)					
Receiving Office	Priority Date Claimed					
UK PATENT OFFICE	13 February 1990 (13.02.90)					
Applicant (Name)						
AMERSHAM INTERNATIONAL PLC ET AL						
BASIS O	F REPORT					
AMENDMENTS AND/OR RECTIFICATIONS ¹ — The amendments Authority in respect of the claims, the description, and/or drawings in	and/or rectifications made before this International Preliminary Examining the above-identified international application are annexed to this report.					
a. [X This report has been established on the basis of the followin	g application documents:					
(X) the application documents as filed	•					
description, pages	as originally filed					
description, pages	filed with your letter of					
description, pages	filed with your letter of					
description, pages	filed with your letter of					
claim(s)	as originally filed					
claim(s)	filed with your letter of					
claim(s)	filed with your letter of					
claim(s)	filed with your letter of					
drawings, sheet/fig.	as originally filed					
drawings, sheet/fig.	filed with your letter of					
D. The amendments resulted in the cancellation of the following she	vets:					
This report has been established as if the amendments indicated of have been considered to go beyond the disclosure as filed.	on the extra sheet have not been made, since, for the reasons indicated, they					
2. PRIORITY ²						
This report has been established as if no priority has been requested:	claimed due to the failure to furnish within the prescribed time limit the					
copy of the earlier application whose priority has been c	laimed.					
translation of the earlier application whose priority has b	een claimed.					
b. This report has been established as if no priority has been	claimed due to the fact that the priority claim has been found invalid					
Thus, for the purposes of this report, the international filling date in	dicated above is considered to be the relevant date.					

BASIS OF REPORT (Continued)
3. UNITY OF INVENTION 3 — The international application does not comply with the requirement of unity of invention.
a. In response to an invitation to restrict or pay additional fees the applicant has:
restricted the claims.
paid additional fees. paid additional fees under protest. Where requested by the applicant, the text of the protest together with the decision
taken thereon are annexed to this report.
neither restricted nor paid additional fees.
b. No invitation has been issued. The opinion of this International Preliminary Examining Authority is that the international application does not comply with the requirement of unity of invention for the following reasons. (specify)
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c. Consequently, the following parts of the international application were the subject of international preliminary examination in
establishing this report:
the parts relating to the restricted claims, that is claims Nos.
the parts relating to the main invention, that is claims Nos
4. NON-ESTABLISHMENT OF REPORT ON QUESTIONS OF NOVELTY, INVENTIVE STEP OR INDUSTRIAL APPLICABILITY 4
The questions of whether the claimed invention appears to be novel, to involve an inventive step or to be industrially applicable have not for the reasons indicated been gone into in respect of:
a. the entire international application
b. claims Nos
for the following reasons:
Said international application, or said claims Nos relate to the following subject matter which does not
require an international preliminary examination. (specify)
The description, claims, or drawings (indicate particular elements) or said claims Nos are so unclear that no meaningful opinion could be formed.
The claims, or said claims Nos are so inadequately supported by the description that no meaningful opinion could be formed.
Said claims Nos are dependent claims and are not drafted in accordance with the second and third sentences of PCT Rule 6.4(a).

CLASSIFICATION OF SUBJECT MATTER (If several classification symbols apply, indicate all.) 5

According to International Patent Classification (IPC) or to both National Classification and IPC

 IPC^{5} B03C 1/00: C07K 3/24: 3/28: C12N 1/02. 7/02. 5/00

REASONED STATEMENT AS TO CLAIMS MEETING CRITERIA OF NOVELTY (N), INVENTIVE STEP (IS) AND INDUSTRIAL APPLICABILITY (IA) * AND CITATIONS * AND EXPLANATIONS * SUPPORTING SUCH STATEMENT

CLAIM	STATEMENT	CITATIONS AND EXPLANATIONS
CLAIM NUMBER	STATEMENT (CRITERIA)	CHATIONS AND EXPLANATIONS
1-12	YES (N.	All claims meet the requirements of industrial applicabilit
_	IS. IA)	novelty and inventive step.
		They are distinguished from the documents cited by the International Searching Authority in that there is no disclosure of the suspension of magnetic particles in a polymer solution prior to precipitating the polymer out of the solution.
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	NON-WRITTEN	DISCLOSURES .	
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	ERTAIN DEFECTS IN THE IN	TERNATIONAL APPL	ICATION 11
	n or contents of the international a		
The claims	are not in two part	t form as requi	red by Rule 6.3(b).
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CERT	AIN OBSERVATIONS ON TI	HE INTERNATIONAL	APPLICATION 12
he following observations on the time of the description have been no	ne clarity of the claims, description ted.	, and drawings or on the q	uestion whether the claims are fully support
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The content	s of the Agent's la	etter dated 10	September 1991 have
been consid			•
	CERT	IFICATION	
Date Demand Submitted		Date of Completion Report	of the International Preliminary Examination
10 Septembe	er 1991 (10.09.01)	· 1	ry 1992 (12 .02.92)
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International Preliminary Exam	ming Authority	Signature of Author	ized Officer
THE PAT	TENT OFFICE		
	RD., NEWPORT	J L FRE	JEMAN / Wheel
GWENT	NP9 1RH	:	- / / ` '

	CUMENTS CONSIDERED TO BE RELEVANT (CONTINUED FROM THE SECOND SHEET	Relevant to Claim No.
ategory *	Citation of Document, 11 with Indication, where appropriate, of the relevant passages	NAMES OF THE PARTY
A	WO, A, 8401503 (COULTER et al.) 26 April 1984 see the whole document; especially page 7, lines 1-5	1-12
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INTERNATIONAL SEARCH REPORT

International Application No PCT/GB 91/00212

A CLASCIFICATION OF CHOISET MATTER (if square) elegification symbols apply indicate all)																			
CLASSIFICATION OF SUBJECT MATTER (if several classification symbols apply, indicate all) According to International Patent Classification (IPC) or to both National Classification and IPC																			
IPC ⁵ :	B 0	3 C	1/	00,	C	07	K	3 /	24,	, 3	/28,	C	12	N	1/02	,	7/0	2,	5/00
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US-A- 3470067	30-09-69	None	
BE-A- 686243	28-02-67	None	
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EP-A- 0281390	07-09-88	AU-A- 1426988 JP-T- 1502319 WO-A- 8806633	26-09-88 17-08-89 07-09-88

YAR J. C. CANARA

(30) Priority data:



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 5: (11) International Publication Number: WO 91/12079 B03C 1/00, C07K 3/24, 3/28 A1 (43) International Publication Date: 22 August 1991 (22.08.91) C12N 1/02, 7/02, 5/00

PCT/GB91/00212 (21) International Application Number:

(22) International Filing Date: 13 February 1991 (13.02.91)

9003253.3 13 February 1990 (13.02.90)

(71) Applicant (for all designated States except US): AMER-SHAM INTERNATIONAL PLC [GB/GB]; Amersham Place, Little Chalfont, Buckinghamshire HP7 9NA (GB).

(72) Inventor; and (75) Inventor/Applicant (for US only): REEVE, Michael, Alan [GB/GB]; 149 Grays Road, Henley-on-Thames RG9 ITE (GB).

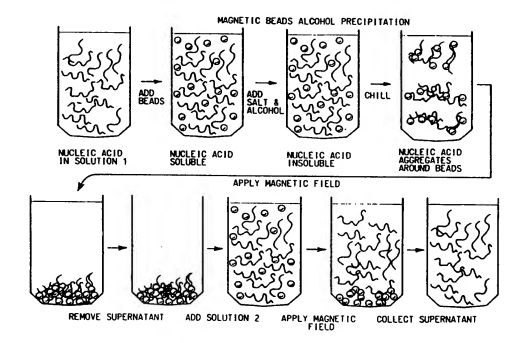
(74) Agent: PENNANT, Pyers; Stevens, Hewlett & Perkins, 2 Serjeants' Inn, Fleet Street, London EC4Y ILL (GB).

(81) Designated States: AT (European patent), BE (European patent), CA, CH (European patent), DE (European patent), DK (European patent), ES (European patent), FR (European patent), GB (European patent), GR (European patent), IT (European patent), JP, LU (European patent), NL (European patent), SE (European patent),

Published

With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments.

(54) Title: METHOD TO ISOLATE MACROMOLECULES USING MAGNETICALLY ATTRACTABLE BEADS WHICH DO NOT SPECIFICALLY BIND THE MACROMOLECULES



(57) Abstract

A method of recovering a biopolymer from solution involves the use of magnetically attractable beads which do not specifically bind the polymer. The beads are suspended in the solution. Then the polymer is precipitated out of solution and becomes non-specifically associated with the beads. When the beads are magnetically drawn down, the polymer is drawn down with them. The polymer can subsequently be resolubilised and separated from the beads.



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DE	Germany	LU	Luxembourg	US	United States of America
DK	Denmark	MC	Monaco		

_ 1 _

Method to isolate macromolecules using magnetically attractable beads which do not specifically bind the macromolecules.

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15

I Introduction

Many techniques in Molecular Biology,
Biochemistry and Chemistry rely upon the process of
precipitation. There are two types of precipitation.

In the first type of precipitation, the components from a complex solution that are not of interest are selectively precipitated. The precipitate and supernatant are then separated (usually by centrifugation or filtration) and the supernatant is kept for further use.

In the second type of precipitation, the components of interest from a complex solution are selectively precipitated. The precipitate and supernatant are separated (again by centrifugation or filtration) and the precipitate is kept for further use. This precipitate may well be redissolved for further use.

Examples of precipitation that are of particular relevance to this invention will now be discussed.

a. Alcohol Precipitation of Nucleic Acid Molecules from Solution:

Alcohol precipitation of nucleic acid molecules from solution is a standard procedure for the concentration and/or purification of these species from complex solutions. Typical methods involve the addition of salt (e.g. 0.1 volumes of 2.5 M sodium acetate (pH 5.2)) to a solution containing nucleic acids followed by addition of an alcohol (e.g. 2.5

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volumes of ethanol). The nucleic acids then precipitate. The precipitated nucleic acid molecules aggregate (usually with the aid of reduced temperatures; e.g. 5 minutes on dry ice) and are 5 recovered by centrifugation. After removal of the supernatant, the pelleted precipitate is normally redissolved in the required volume of an appropriate The nucleic acid may be DNA (partially or wholly single or double stranded), RNA (partially or wholly single or double stranded), mixtures of any of 10 the above or a hybrid RNA/DNA species. The salt used may be sodium acetate, sodium chloride, potassium acetate, potassium chloride, ammonium acetate, ammonium chloride, guanidinium thiocyanate, guanidinium isothiocyanate, guanidinium chloride or mixtures of the above. The alcohol used is normally ethanol or isopropanol.

b. Precipitation of Bacteriophage and Other Viruses from Solution:

Precipitation of bacteriophage and other viruses from solution by the addition of solutions containing high concentrations of highly hydratable polymers, such as polyethylene glycol (PEG), and salts, such as sodium chloride, is a standard procedure for 25 the concentration and/or purification of these species from complex solutions. The bacteriophage or other viruses precipitated in this way may be used for nucleic acid extraction, protein extraction, infection of host cells, structural studies or immunological 30 studies. A typical procedure involves the addition of 0.2 volumes of 20% (w/v) PEG in 2.5 M sodium chloride to the complex solution known to contain the bacteriophage or other viruses. The bacteriophage or other viruses precipitate. The precipitated particles 35

then aggregate (normally with the aid of incubation at reduced temperatures; e.g. 60 minutes at 4°C) and are recovered by centrifugation. After removal of the supernatant, the pellet (comprising precipitated particles of bacteriophage or other viruses) is normally redissolved in the required volume of an appropriate buffer. The bacteriophage may be filamentous (e.g. M13) or complex (e.g. lambda). They may infect bacteria, animal or plant cells and they may be DNA-containing or RNA-containing.

c. Removal of Bacterial DNA. Proteins and Membranes from Bacterial Lysates:

Another type of precipitation of interest to 15 Molecular Biologists is used for the removal of bacterial DNA, proteins and membranes from bacterial lysates containing, in addition to the above, RNA and plasmid DNA and/or cosmid DNA and/or bacteriophage DNA. This forms the basis of the alkaline lysis procedure for preparations of low molecular weight DNA. In this procedure, the bacterial cells (e.g. E.coli) are lysed by treatment with sodium hydroxide (e.g. 200 mM) and the detergent sodium dodecyl sulphate (SDS) (e.g. 0.3-1.0% (w/v)). Addition of a mixture of either sodium or potassium acetate at low pH (e.g. 0.5 times the volume of lysis buffer of 3 M sodium or potassium acetate adjusted to pH 4.8 with acetic acid) leads to the formation of a precipitate containing protein, membrane fragments and the entrapped bacterial DNA. low molecular weight DNA species are not entrapped in 30 this precipitate and can be recovered from the supernatant after centrifugation or filtration of the precipitate. The low molecular weight DNA species can be purified and/or concentrated, along with cellular RNA, by subsequent alcohol precipitation from this supernatant as described above. The DNA species

extracted by this procedure may be plasmid, cosmid or bacteriophage-derived. The volume of cells lysed can be as little as a few microlitres or as large as many litres of bacterial culture.

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II Description of the Invention

In one aspect this invention provides a method of treating a solution of a polymer by the use of magnetically attractable beads which do not specifically bind the polymer, comprising the steps of:

- suspending the magnetically attractable beads in the solution.
- precipitating the polymer out of solution whereby it becomes non-specifically associated with the beads,
 - applying a magnetic field to draw down a precipitate of the beads and the associated polymer, and
- separating the precipitate from a 20 supernatant liquid.

The key to the invention is the use of magnetically attractable beads (hereinafter magnetic beads). The nature of the magnetic beads is not critical, and commercially available beads may be used.

The beads typically have an average diameter in the range 1 to 100 μm , and comprise finely divided magnetizable material encapsulated in organic polymer.

Or the organic polymer may be omitted. Beads of magnetic iron oxide are commercially available.

Such beads have been successfully used in this invention in sizes ranging from below 1 μm up to 40 μm . Even the larger beads remain in suspension at least for the duration of the precipitation step; their subsequent tendency to settle out assists the magnetic field in drawing down the precipitate.

To improve recovery of precipitated polymers,

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the beads may be pretreated to reduce any unwanted tendency to bind the polymers permanently. For example, when the polymers are nucleic acids, the beads may be pre-treated with a phosphate solution. This treatment is believed to phosphatize any exposed magnetisable material, and may not be necessary if the magnetisable material is completely encapsulated in inert polymer.

The beads are preferably added to the solution either before, or together with, a reagent used to precipitate the polymer. Alternatively, the beads may be added after the precipitation step, under conditions to cause the pre-existing precipitate to become associated with them.

The starting solution is preferably aqueous. However starting solutions in polar or non-polar organic solvents are envisaged, particularly when the polymer is of synthetic origin.

generally, it is of particular importance in relation to biopolymers. Biopolymers are polymers found in biological systems. The nature of the biopolymer is not critical to the invention. Biopolymers include nucleic acids (DNA and RNA), proteins, polypeptides, polysaccharides, cell membrane material, bacteriophages, virus, and procaryotic and eucaryotic cells.

At the outset, the polymer or biopolymer is present in solution, the term solution being used broadly to cover permanently stable suspensions in which the polymer molecules are not aggregated.

It is a feature of the invention that the magnetic beads do not specifically bind the polymer. By this feature, the present invention is distinguished from many prior techniques which involve providing a coating on the surface of magnetic beads designed to

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specifically bind the substance to be drawn down out of solution. When the polymer is precipitated out of solution in the presence of the suspended magnetic beads, it becomes non-specifically associated with the beads. When the beads are drawn down by an applied magnetic field, the associated precipitated polymer is drawn down with them. But when in solution, the polymer does not become associated with the beads.

when the solute is of more interest than the solvent, the method may be used either to concentrate an initially dilute solution, or to recover one or more polymers from a mixture of polymers, or for both these purposes in sequence. For a sequence of manipulations, the same beads can conveniently be used. The nature of the liquids used to dissolve or re-dissolve the polymer, and of reagents used to precipitate polymer, are not material to this invention. A skilled reader will have no difficulty in choosing liquids and reagents appropriate to his needs.

In another aspect, the invention provides an automated device for performing this method, which device comprises an automated pipettor and a magnet which may be a permanent magnet or an electromagnet.

The invention will now be discussed with reference to the three types of precipitation given in the introduction.

a. Alcohol Precipitation of Nucleic Acid Molecules from Solution:

Magnetic bead induced precipitate separation can be used to greatly improve the process of alcohol precipitation of nucleic acids. The alcohol precipitation procedure as modified by this invention is shown in Figure 1a. Magnetic beads are added to the nucleic acid in solution. Salt is then added (the magnetic beads can also be added at the same time as

the salt). The nucleic acid is still soluble at this stage. Alcohol is then added. This causes the nucleic acid to come out of solution. The precipitated nucleic acid aggregates around the suspended magnetic beads (which may well act as nucleation sites for this aggregation process). The aggregation stage may be assisted for some types of precipitations by chilling (though chilling does not appear to be necessary for simple precautions of plasmid, phage DNA, RNA and genomic DNA by this method). A magnetic field is then 10 applied to the precipitation. This magnetic field is used to draw the complex of magnetic beads and precipitated nucleic acid to the bottom (or side) of the tube. The supernatant is then removed from the tube. At this point, the precipitate can be washed .15 with ethanol, and/or isopropanol and/or 70% (v/v) ethanol to remove any residual salt, nucleotides, chemicals or organic solvents remaining from treatments of the nucleic acid prior to the precipitation step. The nucleic acid is insoluble in isopropanol, ethanol 20 and 70% (v/v) ethanol. The nucleic acid therefore remains aggregated around the magnetic beads during washing. The washing step can thus be performed vigorously (e.g. by vortex mixing) without risk of losing the precipitate. After the washing step, if 25 performed, the precipitate is redissolved in the required volume of an appropriate buffer in the absence of the magnetic field. Reapplication of the magnetic field to the tube results in just the magnetic beads being drawn to the bottom (or side) of the tube (as the 30 nucleic acid is now dissolved rather than a precipitate as before). The redissolved nucleic acid can now be separated from the magnetic beads by collecting the supernatant containing the dissolved nucleic acid with a pipette whilst the beads are held against the bottom 35 (or side) of the tube by the magnetic field.

The modification of alcohol precipitation by this invention has several clear advantages over the conventional method of precipitation using centrifugation. The procedure, as modified by this invention, is:

- 1. Faster (the modified procedure takes only 1-2 minutes, as opposed to 10-30 minutes for the conventional procedure using centrifugation).
- Not reliant upon centrifugation equipment.
- 10 3. Readily suited to automation (a great many tubes could be placed over a large electromagnet and these could all be alcohol precipitated simultaneously using a multi channel pipetting device).
- 4. Especially effective if the precipitate of nucleic acid is to be washed with isopropanol, ethanol or 70% ethanol (e.g. to remove any residual salt, nucleotides or organic solvents such as phenol). Washing can be performed rapidly with no risk of loss of material as can occur with the conventional method based upon centrifugation (where the pellet often detaches from the bottom of the tube during such washing).

Magnetic bead induced precipitate separation can also be used to greatly improve the process of deproteinization and alcohol precipitation of nucleic 25 acids. The deproteinization and alcohol precipitation procedure as modified by this invention is shown in Figure 1b. DNA is given as the example in Figure 4b, though the process is equally applicable to any type of nucleic acid. Magnetic beads are added to the protein 30 and nucleic acid in solution. Salt is then added (the magnetic beads can also be added at the same time as the salt). The protein and nucleic acid are still soluble at this stage. Alcohol is then added. causes the protein and nucleic acid to come out of 35 The precipitated protein and nucleic acid

aggregate around the suspended magnetic beads (which may well act as nucleation sites for this aggregation process). The aggregation stage may be assisted for some types of precipitations by chilling (though 5 chilling does not appear to be necessary for simple precipitations of plasmid, phage DNA, RNA and genomic DNA with protein extraction by this method). magnetic field is then applied to the precipitation. This magnetic field is used to draw the complex of magnetic beads and precipitated protein and nucleic 10 acid to the bottom (or side) of the tube. supernatant is then removed from the tube. The protein and nucleic acid remain aggregated around the magnetic beads. Phenol and/or phenol/chloroform and/or phenol/ethanol is then added and the magnetic beads 15 resuspended in the absence of the magnetic field. resuspension extracts the precipitated protein from the magnetic beads whilst the nucleic acid remains still attached. A magnetic field is again applied to the tube. This magnetic field is used to draw the complex 20 of precipitated nucleic acid and magnetic beads to the bottom (or side) of the tube. The phenolic supernatant (containing the extracted protein) is then removed from the tube. At this point, the precipitate can be washed 25. with ethanol, and/or isopropanol and/or 70% (v/v)ethanol to remove any residual salt, nucleotides, chemicals or organic solvents remaining. The nucleic acid is insoluble in isopropanol, ethanol and 70% (v/v) The nucleic acid therefore remains aggregated ethanol. around the magnetic beads during washing. The washing 30 step can thus be performed vigorously (e.g. by vortex mixing) without risk of losing the precipitate. the washing step, if performed, the precipitate is redissolved in the required volume of an appropriate buffer in the absence of the magnetic field. 35 Reapplication of the magnetic field to the tube results

in just the magnetic beads being drawn to the bottom (or side) of the tube (as the nucleic acid is now dissolved rather than a precipitate as before). The redissolved nucleic acid can now be separated from the magnetic beads by collecting the supernatant containing the dissolved nucleic acid with a pipette whilst the beads are held against the bottom (or side) of the tube by the magnetic field.

The modification of deproteinization and alcohol precipitation by this invention has several clear advantages over the conventional method of using centrifugation. The procedure, as modified by this invention, is:

- 1. Faster (the modified procedure takes only 5-15 10 minutes, as opposed to 20-40 minutes for the conventional procedure using centrifugation).
 - Not reliant upon centrifugation equipment.
- 3. Readily suited to automation (a great many tubes could be placed over a large electromagnet and these could all be deproteinized and alcohol precipitated simultaneously using a multi channel pipetting device).
- 4. Especially effective if the precipitate of nucleic acid is to be washed with isopropanol, ethanol or 70% ethanol (e.g. to remove any residual salt, nucleotides or organic solvents such as phenol). Washing can be performed rapidly with no risk of loss of material as can occur with the conventional method based upon centrifugation (where the pellet often detaches from the bottom of the tube during such washing).

b. Precipitation of Bacteriophage and Other Viruses from Solution:

35 Magnetic bead induced precipitate separation can be used to greatly improve the process of

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hydrateable polymer/salt precipitation of bacteriophage and other viruses. The hydrateable polymer/salt precipitation procedure as modified by this invention is shown in Figure 2. Magnetic beads, hydrateable polymer (e.g. PEG) and salt (e.g. sodium chloride) are 5 added to the bacteriophage or other viral particles in This causes the particles of bacteriophage or other viruses to come out of solution. precipitated particles aggregate round the suspended. magnetic beads (which may well act as nucleation sites 10 for this aggregation process). The aggregation stage may be assisted for some types of precipitations by chilling (though chilling does not appear to be necessary for simple precipitations of bacteriophage). A magnetic field is then applied to the precipitation. 15 This magnetic field is used to draw the complex of magnetic beads and precipitated particles to the bottom (or side) of the tube. The supernatant is then removed from the tube. The precipitate is redissolved in the required volume of an appropriate buffer in the absence 20 of the magnetic field. Reapplication of the magnetic field to the tube results in just the magnetic beads being drawn to the bottom (or side) of the tube (as the particles of bacteriophage or other viruses are now dissolved rather than a precipitate is before). 25 redissolved particles of bacteriophage or other viruses can now be separated from the magnetic beads by collecting the supernatant containing the dissolved particles with a pipette whilst the beads are held against the bottom (or side) of the tube by the 30 magnetic field.

The modification of hydrateable polymer/salt precipitation by this invention has several clear advantages over the conventional method of precipitation using centrifugation. The procedure, as modified by this invention, is:

- 1. Faster (the modified procedure takes only 1-2 minutes, as opposed to 65-75 minutes for the conventional procedure using centrifugation).
- 2. Not reliant upon centrifugation equipment.
- 5 3. Readily suited to automation (a great many tubes could be placed over a large electromagnet and these could all be hydrateable polymer/salt precipitated simultaneously using a multi channel pipetting device).
- 10 4. Less likely to produce aerosols of bacteriophage and other viruses than the conventional procedure based upon centrifugation. This is safer if the bacteriophage or other viruses are harmful and will result in less airborne microbial contamination in the laboratory.

c. Removal of Bacterial DNA. Proteins and Membranes from Bacterial Lysates:

Magnetic bead induced precipitate separation can also be used to greatly improve the precipitation 20 of bacterial DNA, membranes and proteins from bacterial lysates containing RNA and low molecular weight DNA The preparation of RNA and low molecular weight DNA species as modified by this invention is shown in Figure 3. Bacteria (containing the low 25 molecular weight DNA species of interest) are lysed with a mixture of sodium hydroxide and SDS. releases bacterial DNA, proteins, membranes, RNA and low molecular weight DNA into solution. Magnetic beads and either sodium or potassium acetate are then added 30 This causes the SDS, proteins and membranes at low pH. to precipitate. The precipitate also entraps the bacterial DNA and the magnetic beads. A magnetic field is then applied to the precipitation. This magnetic field is used to draw the complex of magnetic beads and 35 precipitated material to the bottom (or side) of the

tube. The supernatant is then removed from the tube with a pipette whilst the complex of beads and precipitated material is held against the bottom (or side) of the tube by the magnetic field. The low molecular weight DNA can be purified and/or concentrated from this supernatant (along with any remaining cellular RNA that will also be purified) by alcohol precipitation as described above.

The modification of low molecular weight DNA preparation by this invention has several clear advantages over the conventional method of precipitation using centrifugation. The procedure, as modified by this invention, is:

- 1. Faster (the modified procedure takes only 5-10 minutes, as opposed to 30-60 minutes for the conventional procedure using centrifugation).
 - 2. Not reliant upon centrifugation equipment.
- 3. Readily suited to automation (a great many tubes could be placed over a large electromagnet and these could all be precipitated simultaneously using a multi channel pipetting device).

d. DNA Preparation from Bacteriophage or Other Viruses:

has been shown to greatly improve the process of hydrateable polymer/salt precipitation of bacteriophage and other viruses. The hydrateable polymer/salt precipitation has been shown in Figure 2. Magnetic bead induced precipitate separation has also been shown to greatly improve the precipitation of bacterial DNA, membranes and proteins from bacterial lysates containing RNA and low molecular weight DNA species. The preparation of RNA and low molecular weight DNA species as modified by this invention has been shown in Figure 3. The combination of these two procedures can

be used to derive a novel procedure for the purification of low molecular weight DNA from bacteriophage or other viral particles. In this novel procedure, particles of bacteriophage or other viruses are precipitated using the magnetic bead method given in Figure 3. The purified particles are then subjected to lysis by sodium hydroxide and SDS. step separates the coat proteins from the DNA, with both being released into solution. Magnetic beads and either sodium or potassium acetate are then added at This causes the SDS and coat proteins to precipitate. The precipitate also entraps the magnetic A magnetic field is then applied to the This magnetic field is used to draw the precipitation. complex of magnetic beads and precipitated material to 15 the bottom (or side) of the tube. The supernatant is then removed from the tube with a pipette whilst the complex of beads and precipitated material is held against the bottom (or side) of the tube by the The low molecular weight DNA can be magnetic field. 20 purified and/or concentrated from this supernatant by alcohol precipitation as described above. modification of low molecular weight DNA preparation from bacteriophage or other viruses by this invention has several clear advantages over the conventional 25 method of precipitation using centrifugation and other methods. The procedure, as modified by this invention, is:

- Faster (the modified procedure takes only 5 minutes, as opposed to 2-3 hours for the conventional procedure using centrifugation).
 - 2. Not reliant upon centrifugation equipment.
- 3. Readily suited to automation (a great many tubes could be placed over a large electromagnet and these could all be precipitated simultaneously using a multi channel pipetting device).

- 4. Not reliant upon organic solvent extraction (e.g. by phenol).
- 5. Less likely to produce aerosols of bacteriophage and other viruses than the conventional procedure based upon centrifugation. This is safer if the bacteriophage or other viruses are harmful and will result in less airborne microbial contamination in the laboratory.
- 6. Especially effective if the precipitate of nucleic acid is to be washed with isopropanol, ethanol or 70% ethanol. Washing can be performed rapidly with no risk of loss of material as can occur with the conventional method based upon centrifugation (where the pellet often detaches from the bottom of the tube during such washing).

e. Precipitation of Bacteria from Solution

Magnetic bead induced precipitate separation can also be used to effect a novel process of alcohol precipitation of cells e.g. bacterial cells. Magnetic 20 beads are added to the bacteria in solution. then added (the magnetic beads can also be added at the same time as the salt). The bacteria are still soluble at this stage. Alcohol is then added. causes the bacteria to come out of solution. 25 precipitated bacteria aggregate around the suspended magnetic beads (which may well act as nucleation sites for this aggregation process). A magnetic field is then applied to the precipitation. This magnetic field is used to draw the complex of magnetic beads and 30 precipitated bacteria to the bottom (or side) of the The supernatant is then removed from the tube. The precipitate is dissolved in the required volume of an appropriate buffer in the absence of the magnetic Reapplication of the magnetic field to the tube 35 results in just the magnetic beads being drawn to the

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bottom (or side) of the tube (as the bacteria are now dissolved rather than a precipitate as before). The redissolved bacteria can now be separated from the magnetic beads by collecting the supernatant containing the dissolved bacteria with a pipette whilst the beads are held against the bottom (or side) of the tube by the magnetic field.

Alternatively the bacteria can be lysed directly on the beads as described in IIIc for DNA preparation. The procedure as effected by this invention is:

- 1. Faster (the modified procedure takes only 1-2 minutes, as opposed to 5-15 minutes for the conventional procedure using centrifugation).
- 15 2. Not reliant upon centrifugation equipment.
 - Readily suited for automation (a great many culture tubes could be placed over a large electromagnet and these could all be alcohol precipitated simultaneously using a multi channel pipetting device).

III Reduction of the Invention to Practice:

The magnetic beads used were cellulose/ferric oxide (50/50), with a particle size of 1-10 microns diameter. Beads were pretreated by soaking in 100 mM tetrasodium pyrophosphate solution, and stored at 4 degrees in 0.1% (w/v) sodium azide at a concentration of 50 mg/ml.

30 a. An Example of Alcohol Precipitation of Nucleic Acid
Using Magnetic Bead Induced Precipitate Separation:

Example 1

Precipitations of plasmid (e.g. pBR322) can be performed according to the following protocol: Take pBR322 DNA in, for example, 100 µl of TE buffer (10 mM

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Tris-HCl (pH 8.0), 1 mM EDTA (pH 8.0)); add 1/10th volume (i.e. 10 µl) of a solution containing magnetic beads at 50 mg/ml and tRNA at 1 mg/ml (as carrier, this may be omitted if the concentration of nucleic acid is greater than about 10-25 $\mu g/ml$) in 2.5 M sodium acetate adjusted to pH 5.2 with acetic acid; mix; add 2.5 volumes (i.e. 250 µl) of ethanol; mix; place over a permanent magnet to bring down the precipitate; the supernatant; wash the pellet in, for example, 100 µl of 70% (v/v) ethanol by resuspending with a pipette in the absence of the magnetic field, finally removing the supernatant (wash solution) with a pipette in the presence of the magnetic field; redissolve the pellet in the required volume of TE buffer.

No loss occurs on omission of the chilling step for pBR322 DNA. Also, no loss occurs from washing the precipitate with 70% (v/v) ethanol for pBR322 DNA. The above procedure works equally well for human genomic DNA and for RNA.

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Example 2

Precipitation of pBR322 plasmid DNA with deproteinization can be performed according to the following protocol: Take, for example, pBR322 DNA in 20 µl of TE buffer (10 mM Tris-HCl (pH 8.0), 1 mM EDTA (pH 8.0)) containing protein (e.g. a 1/4 dilution of Rainbow Markers (Amersham International)); 1/10th volume (i.e. 2 μ l) of a solution containing magnetic beads at 50 mg/ml and tRNA at 1 mg/ml (as carrier, this may be omitted if the concentration of nucleic acid is greater than about $10-25 \mu g/ml$) in 2.5 M sodium acetate adjusted to pH 5.2 with acetic add 2.5 volumes (i.e. 50 µl) of ethanol; mix; place over a permanent magnet to bring down the precipitate; remove the supernatant; wash the pellet in, for example, 50 µl of phenol (or greater than 60%

(v/v) phenol in ethanol) by resuspending with a pipette in the absence of the magnetic field, finally removing the supernatant (wash solution) with a pipette in the presence of the magnetic field; redissolve the pellet in the required volume of TE buffer. The yield of DNA falls off with less than 40% (v/v) phenol in ethanol used for protein extraction. No losses are incurred with this additional step of protein extraction compared to a protein-free ethanol precipitation.

Ethanol precipitation from a solution heavily contaminated with protein is also seen to be dependent upon the extraction of the contaminating protein by a phenol containing solution (i.e. the DNA cannot be redissolved from the beads if protein extraction has not been performed). The successful extraction of the protein into the phenolic layer by this procedure can clearly be seen when using coloured proteins. The above procedure works equally well for human genomic DNA and for RNA.

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b. An Example of Hydrateable Polymer/Salt Precipitation of Bacteriophage Using Magnetic Bead Induced Precipitate Separation:

25 Example 3

Precipitations can be performed on, for example, 1 ml samples of M13mp8 bacteriophage in 2xTY broth (precleared of bacteria by centrifugation) according to the following protocol: Add 0.4 volumes (i.e. 400 μ l) of 2 mg/ml magnetic beads in 20% (w/v) PEG, 2.5 M NaCl; mix; bring down magnetic beads and precipitate using a permanent magnet; redissolve magnetic bead pellet in the required volume of TE buffer.

At 0.4 volumes, the amount of bacteriophage not brought down by the beads is negligible.

Example 4

DNA preparations can be performed on, for example, 1 ml samples of M13mp8 bacteriophage in 2xTY broth (precleared of bacteria by centrifugation) 5 according to the following protocol: Add 0.4 volumes (i.e. 400 μ l) of 2 mg/ml magnetic beads in 20% (w/v) PEG, 2.5 M NaCl; mix; bring down magnetic beads and precipitate using a permanent magnet; redissolve magnetic bead pellet in 1/5th volume (i.e. 200 µl) of TE buffer; extract with an equal volume (i.e. 200 µl) 10 of phenol; remove aqueous (top) layer; add 1/10th volume (i.e. 20 µl) of a solution containing magnetic beads at 50 mg/ml and tRNA at 1 mg/ml (as carrier, this may be omitted if the concentration of nucleic acid is greater than about 10-25 μ g/ml) in 2.5 M sodium acetate adjusted to pH 5.2 with acetic acid; mix; add 2.5 volumes (i.e. 500 µl) of ethanol; mix; place over a permanent magnet to bring down the precipitate; remove the supernatant; wash the pellet in, for example, 100 μ l of 70% (v/v) ethanol by resuspending with a pipette in the absence of the magnetic field, finally removing the supernatant (wash solution) with a pipette in the presence of the magnetic field; redissolve DNA in the required volume of TE buffer.

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Example 5

DNA preparations can also be performed on, for example, 1 ml samples of M13mp8 bacteriophage in 2xTY broth (precleared of bacteria by centrifugation) according to the following protocol: Add 0.4 volumes (i.e. 400 μ l) of 2 mg/ml magnetic beads in 20% (w/v) PEG, 2.5 M NaCl; mix; bring down magnetic beads and precipitate using a permanent magnet; redissolve magnetic bead pellet in 1/10th original volume (i.e. 100 µl) of 4 M sodium perchlorate in TE buffer; now

add 2.5 volumes (i.e. 250 µl) of ethanol; mix; place over a permanent magnet to bring down the precipitate; remove the supernatant; wash the pellet twice in, for example, 100 µl of 70% (v/v) ethanol by resuspending with a pipette in the absence of the magnetic field, finally removing the supernatant (wash solution) with a pipette in the presence of the magnetic field; redissolve DNA in the required volume of TE buffer.

10 c. An Example of the Removal of Bacterial DNA.

Proteins and Membranes from Bacterial Lysates Using
Magnetic Bead Induced Precipitate Separation:

Example 6

15 pUC19 DNA can be extracted from E.coli MC1061 cells by the following protocol: Take, for example, 250 µl of bacterial culture; add 1/5th volume (i.e. 50 μ l) of 1.2 M NaOH, 1.2% (w/v) SDS; mix; minutes at room temperature; now add 3/5th volume (i.e. 150 µl) of 10 mg/ml magnetic beads in 3 M 20 potassium acetate adjusted to pH 4.8 with acetic acid; mix; bring down precipitated material with a permanent magnet and keep supernatant; isopropanol precipitate the supernatant as follows: add 1/10th supernatant volume (i.e. 45 µl) of a solution containing magnetic 25 beads at 50 mg/ml and tRNA at 1 mg/ml (as carrier, this may be omitted if the concentration of nucleic acid is greater than about 10-25 µg/ml) in 2.5 M sodium acetate adjusted to pH 5.2 with acetic acid; mix; supernatant volumes (i.e. 270 µl) of isopropanol; mix; 30 place over a permanent magnet to bring down the precipitate; remove the supernatant; wash the pellet in, for example, 100 μ l of 70% (v/v) ethanol by resuspending with a pipette in the absence of the magnetic field, finally removing the supernatant (wash 35 solution) with a pipette in the presence of the

magnetic field; redissolve DNA in the required volume of TE buffer. Preparations can be incubated with 10 μ g/ml ribonuclease A for 10 minutes at 37°C before analysis.

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Example 7

pUC19 DNA can also be extracted from E.coli MC1061 cells by the following protocol: Take, for example, 500 µl of bacterial culture; add 1 volume (i.e. 500 µl) of a solution containing magnetic beads 10 at 5 mg/ml in 0.2 M sodium acetate (adjusted to pH 5.2 with acetic acid) dissolved in ethanol; mix; down precipitated bacteria with a permanent magnet; remove supernatant and discard; redissolve bacterial pellet in, for example, 300 µl of 0.2 M NaOH, 0.2% (w/v) SDS; mix; incubate 2 minutes at room temperature; now add 1/2 volume (i.e. 150 μ l) of 3 M potassium acetate adjusted to pH 4.8 with acetic acid; mix; bring down precipitated material with a permanent magnet and keep supernatant; isopropanol precipitate 20 the supernatant as follows: add 1/10th supernatant volume (i.e. 45 µl) of a solution containing magnetic beads at 50 mg/ml and tRNA at 1 mg/ml (as carrier, this may be omitted if the concentration of nucleic acid is greater than about 10-25 $\mu\text{g/ml}$) in 2.5 M sodium acetate 25 adjusted to pH 5.2 with acetic acid; mix; add 0.6 supernatant volumes (i.e. 270 µl) of isopropanol; mix; place over a permanent magnet to bring down the precipitate; remove the supernatant; wash the pellet in, for example, 100 μ l of 70% (v/v) ethanol by 30 resuspending with a pipette in the absence of the magnetic field, finally removing the supernatant (wash solution) with a pipette in the presence of the magnetic field; redissolve DNA in the required volume of TE buffer. Preparations can be incubated with 35 10 μg/ml ribonuclease A for 10 minutes at 37°C before

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analysis.

d. An Example of DNA Extraction from Bacteriophage
Using Magnetic Bead Induced Precipitate Separation
for: Hydrateable Polymer/Salt Precipitation.
Removal of Coat Proteins and Alcohol Precipitation
of the DNA:

Example 8

M13mp8 phage can be precipitated with magnetic beads, PEG and NaCl as described in IIIb. DNA can then be prepared by the alkaline lysis procedure as described in IIIc (dissolving the PEG/NaCl/magnetic beads precipitate of bacteriophage particles in, for example, 250 µl of TE buffer for alkaline lysis). The alkaline lysis method gives M13mp8 DNA at about half the yield of the phenol extraction preparation.

IV Other Types of Precipitation:

These include the following:

Precipitations of bacteria, tissue culture cells and blood cells by suitable precipitants (e.g. an equal volume of ethanolic 0.2 M sodium acetate adjusted to pH 5.2 with acetic acid for E.coli) and magnetic bead induced precipitate separation.

Ammonium sulphate precipitation of proteins with magnetic bead induced precipitate separation.

Precipitation of proteins by salts other than ammonium sulphate and magnetic bead induced precipitate separation (e.g. sodium perchlorate, sodium iodide, guanidinium chloride, guanidinium thiocyanate, guanidinium isothiocyanate and other chaotropic agents).

Precipitation of proteins by denaturants and magnetic bead induced precipitate separation.

Precipitation of proteins by detergents and

magnetic bead induced precipitate separation.

Precipitation of nucleic acids by the detergent cetyl trimethyl ammonium bromide and magnetic bead induced precipitate separation.

Precipitation or proteins and/or nucleic acids with agents such as trichloroacetic acid (that denature due to extremes of pH) and magnetic bead induced precipitate separation.

Selective RNA precipitations from lithium to chloride and magnetic bead induced precipitate separation.

Selective precipitations of nucleic acids from other nucleic acids (e.g. precipitations of high molecular weight DNA from oligodeoxyribonucleotides and/or deoxynucleotide polyphosphates) which may work better using magnetic bead induced precipitate separation than centrifugation.

Immune precipitations and magnetic bead induced precipitate separation.

20 Complement fixation precipitations and magnetic bead induced precipitate separation.

Blood clotting precipitations and magnetic bead induced precipitate separation.

Latex bead precipitation assays and magnetic bead induced precipitate separation.

Haemagluttination assays and magnetic bead induced precipitate separation.

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CLAIMS

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- 1. A method of treating a solution of a polymer by the use of magnetically attractable beads which do not specifically bind the polymer, comprising the steps of:
- suspending the magnetically attractable beads in the solution,
 - precipitating the polymer out of solution whereby it becomes non-specifically associated with the beads,
- applying a magnetic field to draw down a precipitate of the beads and the associated polymer, and
 - separating the precipitate from a supernatant liquid.
- 20 2. A method as claimed in Claim 1, comprising the additional steps of:
 - adding liquid to the precipitate to redissolve the polymer and re-suspend the beads.
 - applying a magnetic field to draw down the
- 25 beads, and
 - separating a supernatant liquid containing the polymer from the beads.
 - 3. A method as claimed in Claim 1 or Claim 2, wherein the solution is in an aqueous medium.
- 30 4. A method as claimed in any one of Claims 1 to
 - 3, wherein the polymer is a biopolymer.
 - 5. A method as claimed in Claim 1 or Claim 2, wherein the biopolymer is nucleic acid.
 - 6. A method as claimed in Claim 4, wherein the
- 35 biopolymer precipitated comprises protein as well as nucleic acid.

- 7. A method as claimed in Claim 6, comprising the additional steps of:
- adding liquid to the precipitate to selectively re-dissolve the protein and re-suspend the beads,
 - applying a magnetic field to draw down a precipitate of the beads and the associated nucleic acid,
- separating a supernatant liquid containing 10 the protein from the precipitate,
 - adding liquid to the precipitate to redissolve the nucleic acid and re-suspend the beads,
 - applying a magnetic field to draw down the beads, and
- separating a supernatant liquid containing the nucleic acid from the beads.
 - 8. A method as claimed in Claim 4, wherein the biopolymer is bacteriophage and/or virus and/or cell.
- 9. A method as claimed in Claim 4, wherein the starting solution comprises a mixture of similar biopolymers, one of which is selectively precipitated out of solution in the presence of the beads.
 - 10. A method as claimed in Claim 9, wherein the starting solution is a cell lysate comprising protein,
- membrane, bacterial DNA and low molecular weight nucleic acids, and the biopolymer precipitated out of solution comprises the protein, membrane and bacterial DNA but not the low molecular weight nucleic acids.
- 11. A method for recovering low molecular weight nucleic acids from a starting solution of bacteriophage and/or virus, which method comprises the steps:-
 - precipitating the bacteriophage and/or virus and/or cell by the method of Claim 8,
- lysing the bacteriophage and/or virus to
 form a cell lysate solution, and
 - treating the cell lysate solution by the

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method of Claim 10.

12. An automated device for performing the method of any one of Claims 1 to 11, which device comprises an automated pipettor and a magnet.

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